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# MODELOCKING AND WAVEGUIDE AMPLIFIERS USING Cr: FORSTERITE AND Cr: YAG

**Cornell University** 

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# Final Report

## Contract F30602-94-C-0030

# Modelocking and Waveguide Amplifiers Using Cr:Forsterite and Cr:YAG

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### Final Report

#### Contract F30602-94-C-0030

Modelocking and Waveguide Amplifiers Using Cr:Forsterite and Cr:YAG

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#### 1. Introduction

Two initiatives were begun in this effort. First, we attempted to transfer the technology of modelocking a Cr:YAG laser to the Rome Labs team, and second, we began investigating new forms of waveguide amplifiers based on index matched nanocrystals of Cr:forsterite and Cr:YAG.

In the modelocking effort, we successfully demonstrated the first Kerr-lens modelocking of a Cr:YAG laser in our lab at Cornell. Over the period of this contract we worked with the staff at Rome Labs to transfer the techniques for modelocking to their laser. We interacted with Steve Johns, Mark Krol, and Mike Hayduk at Rome Labs in the construction of a similar laser. Pollock made several trips to Rome Labs during the year to help get the Cr:YAG laser operating in the cw mode. While there, continuous wave powers of over one half watt of power were developed from the Cr:YAG laser, and subsequently the Rome Lab team was able to increase the power to over one watt. Following the methods described by Pollock's graduate students and their publications, the Rome Labs members have attempted to modelock the Cr:YAG laser, but have as yet had no success. Steve Johns and Mike Hayduk came to Cornell in April 1994 to see the modelocked laser and to discuss with graduate student Alphan Sennaroglu different techniques for modelocking.

In the thin film amplifier work, graduate student Duane Barber has been investigating the emission properties of thin films made from powder composites of Cr:forsterite, and Cr:YAG. The composite films consist of nanocrystals of Cr-doped particles suspended in a refractive index-matching polymer, spun onto a glass slide. The film thicknesses are one the order of one micron. Even with such a thin sample, Barber has observed significant fluorescent emission from the films, and can characterize their emission lifetime as well. The passive loss of these waveguides was determined to be approximately 6 dB/cm.

#### 2.0 Laser Research

A schematic of Cr:YAG laser cavity is shown below in Fig. 1. The crystal is located at the focus of a z-folded optical resonator. The optical cavity design is relatively straightforward. The critical aspects of the laser are the temperature control of the crystal, and

the optical system used to couple the pump beam into the laser crystal. The crystal in all systems we worked with was mounted on a thermoelectric cooler so that the temperature could be actively maintained below room temperature. We found performance improved with both host materials (forsterite and YAG) as the temperature decreased, but that below about 3 C the optical surfaces became fogged with condensed water from the air. We compromised between cooling the crystal for better power and yet not fogging the crystal surface by operating with crystal temperatures in the 8-12 °C range.

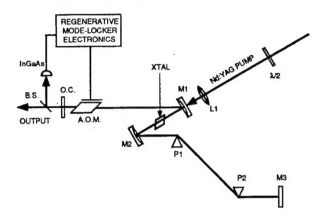


Fig. 1. Schematic of the modelocked solid state laser.

### 2.1 Thermal transfer improvements

One of our key discoveries we made this year was that the transverse size of the laser crystal could be reduced, with a dramatic increase in the laser output power. This was discovered by accident. We inadvertently damaged the face of a Cr:forsterite crystal with dimensions  $5 \times 5 \times 20$  mm. We sliced the crystal into four pieces, each  $2 \times 2 \times 20$  mm, and found that the damage was localized to only one piece. Using one of the remaining sound crystal pieces, we constructed a laser, and found the output power to be almost double that of previous results. We believe the reason for this is due to the reduced path that heat must flow to leave the crystal. The total thermal resistance between the pumped region of the crystal and the copper heat sink is reduced by approximately a factor of four with the smaller dimensions. Therefore the pumped region can operate at a cooler temperature while being pumped at the same powers we used for the large crystal. A second benefit of this discovery is that we can now make several lasers from one purchased crystal, and get better performance at the same time. While our initial results were obtained from using forsterite, they should be directly transferable to the Cr:YAG system. We are pursuing that at the present time.

A recent result we are just beginning to test is the use of indium solder to create the thermal bond between the copper heat sink and the laser crystal. Typically we have placed 0.005" indium foil between the crystal and copper heat sink to act as a compliant thermal transfer medium to allow for better contact between the crystal and heat sink, and to provide some ability for the crystal to expand without danger of cracking. The problem with the indium foil is that it tends to form an oxide on the surface, and the bond between the crystal and indium is not intimate. This dramatically increases the thermal resistance of the interface. We have recently mastered the art of soldering the crystal directly to

the copper heat sink using a hot plate, some soldering flux ("Stay-Clean"), and a clean laser crystal. Initial thermal measurements show that the heat transfer is approximately a factor of 2 better than our standard indium foil technique. Figure 2 shows preliminary data on the crystal cooling efficiency. We are in the process of testing this method for laser operation. If it works as well as anticipated, we may see a further increase in laser performance when operated as before (at 5 C), or we may find that we can eliminate the need to a TE cooler on the laser. Either result would be an advance.

#### 

Fig. 2. Thermal energy transfer from a forsterite crystal using four different bonding techniques. The soldering technique is dramatically better for heat removal.

0.95

1.05

0.85

# 2.2 Cr:YAG Modelocking Attempts

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0.75

We had found in previous work that modelocking the Cr:YAG laser was impossible when we operated at a wavelength near the peak of the output power curve for the laser (1.45  $\mu$ m), which happens to coincide with the strong atmospheric absorption due to water. The absorption primarily introduces a strong dispersion into the cavity which cannot be compensated or overcome with bulk optics. Modelocking was only possible at wavelengths greater than 1.5  $\mu$ m. However, in interacting with the Rome Labs group, we found that their crystal would not tune out to wavelengths much beyond 1.52  $\mu$ m, which made the prospects for modelocking poor. Several attempts were made to fix this problem, including replacing all the optics in their cavity with mirrors we had successfully used here at Cornell. Subsequent to one of Pollock's visits to Rome Labs, it was determined that the absorption of the Cr:YAG crystal at the pump wavelength was significantly less than that observed by the Cornell group. A new crystal was obtained with specifications that the small signal absorption of the pump power exceed certain values.

The new crystal was tested, and initially there was not much improvement over the original tuning range. However, after much iteration, replacement of mirrors, new tuning elements, etc., the Rome Labs group was able to get the new crystal to tune out to beyond  $1.57\ mu$ m. This problem seems to be solved.

After several trips to Rome by Pollock, there still has been no successful modelocking of the Cr:YAG laser. However, concurrent work on Cr:forsterite at Cornell by graduate student Martin Jaspin finally led to the discovery of a successful, and repeatable, method to modelock these lasers. The technique critically relies on observing the spatial mode structure of the laser. As the spacing between the two curved mirrors is changed, the normal Gaussian shaped mode begins to acquire significant higher-order mode structure. When these higher order spatial modes appear, the laser will generally modelock, with 50 fsec pulses. Fig. 3 shows the amplitude autocorrelation of the Cr:forsterite laser during modelocking, and Figure 4 shows the output spectrum. The output of this laser is very quiet and reliable. We are in the process of transferring this technology to Rome Labs for the Cr:YAG laser research.

### Modelocked Forsterite Autocorrelation

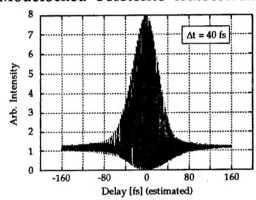


Fig. 3. Autocorrelation of the modelocked output of the Cr:forsterite modelocked laser.

### Modelocked Forsterite Spectrum

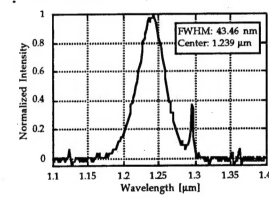


Fig. 4. Spectral content of the modelocked Cr:Forsterite laser. (The spectrum shown here does not correspond to the temporal pulse shown in Fig. 3).

### 3. Thin Film Amplifier Research

Our previous work on thin film amplifiers relied on depositing films of Cr:forsterite on glass using a sol-gel deposition technique which involved spinning on the sol, baking it dry at 600 C, then spinning on another coat, etc. After ten coatings, the film could be

built up to approximately 1  $\mu$ m in thickness, but unfortunately it was heavily fractured due to residual stress caused by the difference in thermal expansion coefficients of the thin film material and the glass substrate. To counter this, we began exploring the use of nanocrystals embedded in an index matching matrix made of polymer or glass. Forsterite particles approximately 50-150 nm in diameter were formed by sintering a sol under the proper pressure and temperature conditions. Powder samples were measured for absorption, emission, and emission lifetime, and were found to be consistent with bulk Cr:forsterite.

Graduate student Duane Barber collaborated with Prof. Chris Ober of the Material Science and Engineering Department, and Prof. Jim Burlitch of Chemistry, to do the index matching of these particles in polymer and glass hosts. The first work was done using nanocrystals of Cr-doped forsterite suspended in a refractive index-matching polymer, and spun onto a glass slide. The films he obtained were approximately  $1\mu$ m thick. Even with such a thin sample, Barber has observed significant fluorescent emission from the films, and can characterize their emission lifetime as well. A sample of the emission obtained from a Cr:forsterite sample is shown in Fig. 5. The data is notable due to the lack of significant emission from  $Cr^{3+}$ , even though green light from an argon ion laser is used to excite the sample. The emission peak shown is characteristic of the  $Cr^{4+}$  transition.

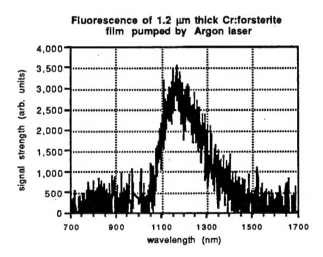


Fig. 5. Spectral emission profile from a Cr:forsterite thin film, pumped by an argon laser

Because forsterite is a biaxial crystal, it is impossible to perfectly index match it in an isotropic medium. Therefore, we can expect that there will be scattering losses associated with the randomly oriented particles in a index-matched host medium. The Rayliegh scattering due to these particles (assuming the particle density is below the percolation point) is given by

 $\frac{P_{scat}}{P_0} = \rho \frac{(n'-n)^2}{n^2} \frac{V^2}{\lambda^4} 24\pi^4$ 

where  $\rho$  is the number of nanocrystals per unit volume, n' is the index of refraction of the scatterers, n is the index of refraction of the host material, V is the volume of a single scatterer, and  $\lambda$  is the wavelength of light. Making the particle size small reduces the

loss due to scattering. Assuming a power gain as high as 50% per cm for forsterite, the trade-off between loss and gain can be evaluated. We have numerically determined that particles on the order of 100-200 nm are the optimum size for this system.

The surfaces of the spun-on polymer films were found to be much too rough to create a low loss waveguide. Losses on the order of 10 dB/cm were measured at a wavelength of 633 nm. Furthermore, once the polymer refractive index matrix had been properly processed to create the nanocrystals, the one micron thickness was too thin to conveniently couple energy into it. With such a high excess loss due to the surface, it is unlikely that we would be able to generate net gain from a thin-film waveguide amplifier. Based on this and discussions with Prof. Chris Ober of the Material Science and Engineering Department, we decided to try building an evanescently coupled amplifier based on nanoparticle composites. A thin film waveguide has been constructed using SiO2/TiO2 glass on a glass waveguide. This waveguide is approximately 1  $\mu$ m thick, and can support one mode. On top of this waveguide, we coated a 10  $\mu m$  thick layer of refractive index matched polymer filled with Cr:forsterite nanoparticles. The evanescent wave of the the guided mode couples to the gain medium, but does not extend far enough into the gain medium to 'see' the rough top surface. Initial measurements of the scattering loss from this structure show that it falls in the 5-8 dB range. We attribute some of this to input scattering at the end-fire input and exit planes of the waveguide. We have not yet been able to identify the source of the rest of the loss. A plot of the loss data, obtained by measuring the scattered light from above the waveguide as it propagates along the waveguide, is shown in Fig. 6.

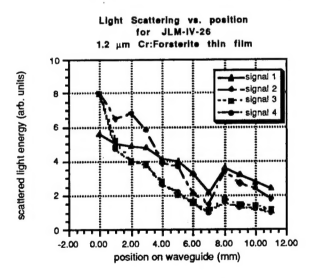


Fig. 5. Plot of the amount of light left in the thin-film waveguide as a function of distance along the waveguide.

#### 4. Conclusion

In this work, we have interacted closely with the group at Rome Labs to get a Cr:YAG laser operational and modelocked. While we have obtained good operating power from this laser, we were unable to successfully transfer the techniques of femtosecond modelocking to the Labs, but we were able to transfer the technology for operating and maintaining

power cw lasers based on Cr:YAG.

The thin film waveguide research demonstrated the concept of using optical nanocomposites in real devices. This has, to our knowledge, never been tried before. Our first results demonstrated that waveguiding can be achieved in nanocomposite materials, and that the losses are not unreasonable for a first effort.

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